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Sustainable platinum recycling through electrochemical dissolution of platinum nanoparticles from fuel cell electrodes

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Abstract

Recycling of the platinum group metals (PGMs) containing industrial wastes obtained from proton exchange membrane fuel cells (PEMFCs), electrolyzers, catalytic convertors and others is of high strategical importance for industrial sustainability. In this work, a highly efficient and environmentally friendly platinum recycling method through potentiodynamic cycling in dilute acidic chloride solutions is demonstrated and optimized to recover platinum from fuel cell electrodes. The process parameters such as upper and lower potential limits are optimized to be 1.6 and 0.4 V vs. RHE. Both the dilute acidic and the acid free Cl− containing electrolytes may be used for the dissolution. Moreover, parameters such as protocol reliability, quantification method and comparison, electrode interface structure, dissolution product and mechanisms etc. are discussed. A study in 1 M HCl shows Pt dissolution rates as high as ~30 µg/cycle for potential cycling between 0.4 and 1.6 V (scan rate: 100 mV/s) for a PEMFC electrode with initial Pt loading of ~470 µg. The approach demonstrates a methodology for fast parameter screening on electrocatalyst dissolution and a proof of concept industrial recycling of spent electrocatalysts.

Keywords: Platinum nanoparticle, Dissolution, Recycling, PEMFC, Electrochemistry, Potentiodynamic
Highlights:

- A recycling oriented Pt nanoparticle electrochemical dissolution protocol is described
- Parameters such as potential limits, electrolyte, concentration etc. are optimized
- More than 95% Pt catalyst can be dissolved using the described process

1. Introduction

Proton exchange membrane fuel cell (PEMFC) using hydrogen as fuel to generate electricity and water represents a fast growing clean and renewable energy conversation technology. Many countries have launched ambitious goals and concrete actions to boost the development and demonstration of the technology: Fuel Cell Projects from Department of Energy (DOE) in U.S., the Fuel Cell Programs from New Energy and Industrial Technology Development Organization (NEDO) in Japan, Fuel cells and Hydrogen Joint Undertaking (FCHJU) in Europe etc.

Among different challenges, cost reduction and material availability, especially platinum group metal (PGM) electrocatalyst, is one of the most acute factors decelerating the ultimate massive commercialization of PEMFCs. Although the required catalyst loading in a PEMFC electrode has reduced by a factor of > 20 in the past decades, still it is too high to be competitive enough.\[^1\] Different strategies have been explored to optimize\[^2-3\], minimize,\[^4\] extent\[^5\] or replace\[^6\] the use of the precious metals as catalysts. However, high performance PEMFCs with PGM-free catalyst still remains a challenge due to change of electrode structures and limited durability from e.g. demetalation and carbon oxidation.\[^7\]

Platinum is the state-of-the-art and most frequently used element as electrocatalyst\[^8\] for PEMFCs as an emerging technology, and in many other well established industries such as automobiles (catalytic convertors), jewelry, chemical synthesis, pharmaceuticals etc., which contribute to a large demand of the element, around 260 tons with 4% steady increase annually.\[^9\] However, platinum is also one of
the least abundant elements in the Earth crust, and more than 88% of the natural reserve localized in South Africa and Russia. These may trigger market supply risk and price instability, if only primary resource is used. Therefore, PEMFC catalyst production based on secondary resource or recycling is highly attractive; moreover, secondary production has various advantageous over primary production, due to the significantly reduced energy consumption and environmental impact. Laforest et al. (2016) carried out a comprehensive environmental assessment on the life-cycle of a PEMFC membrane electrode assembly (MEA). They concluded that more than half of the environmental impact can be avoided if platinum is recovered in the end-of-life MEA.

PGM recycling is a well-established business since the 80’s. A combination of primary and secondary production using pyrometallurgical infrastructure is a common practice, especially for spent auto catalysts. However, pyrometallurgy is not feasible for high volume fuel cell MEA recycling, due to the presence of fluorine compounds. Incineration of the fluoropolymers contained in proton conductors, the hydrophobically treated gas diffusion layer using Teflon®, and the catalyst layer’s binding agent leads to the formation of hydrogen fluoride (HF), a toxic compound that is both harmful to health, environment and highly corrosive for the infrastructure. In addition, through incineration the valuable membrane material, perfluorinated sulfuric acid (PFSA), is essentially destroyed in the process, and therefore a loss of the value. Classic hydrometallurgy has also been practiced for MEA recycling, where high concentration of strong acids and oxidants, e.g. HNO₃, H₂SO₄, HCl or H₂O₂, were used for extraction. However, they are typically unpleasant to work with and have a high requirement for process infrastructure.

In recent years, platinum electrochemical dissolution phenomenon has been intensively studied by several groups, aiming either to understand the degradation mechanisms of Pt electrocatalysts through Pt dissolution under potentiostatic and/or potentiodynamic conditions, which lead to serious
catalyst durability issues, or to develop a process for recovery of PGMs from the end-of-life electrocatalysts,[16, 29-32] Significant understanding of the dissolution mechanisms under different conditions has been developed recently through the studies using a combination of electrochemical potential controlling systems and highly sensitive analytical instrumentation, such as the quartz crystal microbalance (QCM),[33] inductively coupled plasma - mass spectrometry (ICP-MS),[18, 24, 28, 34] X-ray absorption spectroscopy[25], etc. The direct evidence on potentiodynamic dissolution through both anodic and cathodic sweep[21, 25, 27] is one of the most significant contributions in this area. However, the emphasis of most of such pioneering works is the durability of Pt electrocatalysts under fuel cell or electrolyzer related operation conditions,[18, 21, 24-25, 35-36] which does not directly serve the purpose of complete Pt-dissolution or recycling. Nevertheless, comparatively faster degradation of the Pt electrocatalysts in presence of Cl− has motivated to employment of the electrochemical potential cycling to recover Pt from the spent catalysts.[18, 28, 37-38] A few groups[39-40] have further investigated Pt recycling using electrochemical treatment in Cl− containing acidic baths; nevertheless, most of the studies were carried out on bulk material, which limits the possibility for high throughput sampling, requires long treatment time and does not always represent the properties of the corresponding high surface area catalysts, which can be dissolved under comparatively mild conditions (dilute acids, room temperature).

In such a study to recover Pt from nanoparticulate Pt through potentiodynamic treatment, Latsuzbaia et al. (2015) have investigated the influence of Cl− concentration on the dissolution of Pt from a fuel cell electrode by potential cycling between 0.5 and 1.1 V in 0.1 M HCl. In the selected potential window, dissolution through formation of Pt2+ species takes place to dissolve ~0.35 mg Pt on a time scale of 3 – 5 h.[41] However, for large-scale applications, dissolution rates much faster than those reported are desired. Kanamura et al. (2016) have investigated Pt dissolution from MEAs by square wave potential cycling in dilute acids and suggested an optimal upper potential to be 1.4 V for the Pt
dissolution. However, the lower potential was kept constant at 0.1 V. Such a low value of lower potential may not be suitable for efficient Pt dissolution due to the increased probability of redeposition of the dissolved Pt species. Potentiodynamic dissolution through electroless manipulation of the Pt surface potential, having an advantage of its applicability irrespective of the electronic connectivity of the Pt nanoparticles, has been reported by Hodnik et al. (2016), where, high dissolution rates have been achieved by cyclic oxidation and reduction of Pt surface through application of oxidative and reductive gases (e.g. ozone and carbon monoxide). The current work presents feasibility and development on electrochemical platinum catalyst dissolution as a user friendly and environmentally friendly process by using potential control and dilute acidic solutions, to efficiently extract and recover platinum catalyst from its original electrode in form of soluble specious. The process demonstrates a proof-of-concept methodology for platinum group metal recycling and will bring inspiration to an economic, efficient and eco-friendly metallurgical process using electrochemistry as a platform and possibility towards commercialization.

2. Experimental

2.1 Protocol for catalyst electrochemical dissolution

The dissolution behavior of high surface area PEMFC platinum catalyst is highly desirable and valuable information for feasibility assessment and optimization of catalyst recycling. However, platinum nanoparticles usually supported on conductive porous carbon impose several practical challenges for electrochemical dissolution studies, e.g. tedious separation steps between solid and liquid phase, large deviation due to sample loss from reaction or processing, small sample size due to high value of the material etc. Further, for large electrode areas, connection to external power source may be important factor as poor current distribution and potential drops due to Ohmic resistance lead to inhomogeneous dissolution of Pt.
To avoid the aforementioned issues, a user friendly, fast screening protocol for electrochemical dissolution studies on platinum, or in general any electrocatalyst, has been developed. As presence of several metals/alloys and/or support matrices may alter the reaction mechanics leading to low accuracy in reaction kinetic quantification, a relatively simple electrode containing Pt nanoparticles (instead of other typical Pt-alloy electrodes used for PEMFCs) as the catalyst has been studied here. Main features of the procedure consist of:

- Catalyst nanoparticles stabilized in the form of electrode is used for the dissolution study
- Catalyst loading estimation by quantitative analysis, e.g. X-ray fluorescence (XRF) spectroscopy
- Electrode connection with a graphite rod and catalyst dissolution through potentiodynamic treatment (electrochemical dissolution)
- Dissolution element quantification using solution based spectroscopies, e.g. atomic absorption spectroscopy (AAS) or UV-visible spectroscopy.
- Residual element quantification using XRF

Figure 1. Schematic illustration on catalyst electrochemical dissolution evaluation protocol
2.2 Materials and methods

**Materials:** Hydrochloric acid (1 M HCl; Reagent Ph. Eur.; BDH chemicals), Ammonium chloride (NH₄Cl; purity: >99.99%; Reagent Ph. Eur.; BDH chemicals), sulfuric acid (H₂SO₄; EMSURE® grade, assay > 95 – 97%, Merck, Germany), nitric acid (HNO₃; assay: 65%, Sigma-Aldrich®) and Milli-Q water (type I, resistivity ≥18.2 MΩ·cm at 25 °C) were used to prepare different electrolyte solutions. The PEMFC electrodes used in this study were prepared at IRD fuel cells A/S by spray coating of the carbon supported Pt-nanoparticles (HiSPEC 9100 Pt/C catalyst)/Ionomer ink on a gas diffusion layer (GDL; SIGRACET 35DC) to obtain a Pt loading of ~0.6 mg/cm² and an ionomer (Nafion®) content of ~30 wt.%. To investigate the effect of ionomer content on the Pt dissolution, electrodes with Pt loadings of ~0.6 mg/cm² and varying Nafion® contents of 0, 15, 30 and 45 wt.% were prepared at IRD fuel cells A/S by tape casting and spraying of the HiSPEC 9100/Ionomer inks on the SIGRACET 35DC GDL.

**Electrochemical studies:** A three-electrode setup consisting of a working electrode (WE), a counter electrode (CE; Graphitic carbon rod; ~5 mm diameter) and a reference electrode (RE; Hg/Hg₂SO₄; REF 601 Radiometer®) was used for the electrochemical dissolution study. A 10 mm diameter sheet stamped from the catalyst coated GDL was used as the WE, while a graphite rod (0.5 mm diameter) was used as the current collector, as shown in the schematic of Fig. 1a. All potentials have been reported w.r.t. the reversible hydrogen electrode (RHE) by measuring the potential of the Hg/Hg₂SO₄ electrode w.r.t. an RHE in same electrolyte and shifting the measurement potentials accordingly. A Zahner® IM6e electrochemical workstation was used to perform the electrochemical studies in different electrolytes (12 mL; no gas purging; room temperature) and under different potentiodynamic conditions, i.e., potential limits and number of potential cycles (N). The electrolytes were kept
stagnant during the dissolution study, except for the experiment to study the effect of convection flow, where stirring at 750 rpm was applied during the potential cycling.

**Dissolution quantification:** Pt loadings on the WE before and after the electrochemical treatment were determined by using a calibrated Thermo Scientific Niton XL3t GOLDD+ XRF analyzer. The initial and residual Pt loading values ($W_0$ and $W_f$, respectively) were used to determine the $\%$ dissolution of Pt (equation 1) during the electrochemical dissolution.

$$
\%\text{dissolution} = \frac{W_0 - W_f}{W_0} \times 100
$$

(1)

The amount of dissolved Pt in the post-dissolution electrolyte was also measured through atomic absorption spectroscopy (AAS) using a Graphite Furnace Agilent 200 Series AA analyzer. The measurements were performed using a commercial Pt standard (Sigma-Aldrich®) of 250 µg/L and by diluting the electrolytes with a dilution factor (DF) of 200. Moreover, an alternative approach for determination of $\%$ dissolution through UV-vis spectroscopy was demonstrated using a Lambda 900 UV/VIS/NIR spectrometer (Perkin Elmer). UV-Vis spectroscopy was also used to study the chemical state of the dissolved Pt species. The technique is further used to study the Pt-dissolution rates with applied potential during a potential cycle. Electrolyte samples (100 µL) were collected at different potentials (potential step size: 0.2 V) during an electrochemical potential cycle (start: 1.2 V, max.: 1.6 V, min.: 0.4 V and end: 1.6) at a scan rate of 10 mV/s in 0.1 M HCl and 0.1 M HCl+0.1 M CuCl$_2$ electrolytes. The comparatively slow scan rate (10 mV/s) was used to enable the sample collection at a certain potential, while magnetic stirring at 1000 rpm was used to ensure homogeneous distribution of the Pt species in the electrolyte. UV-Vis spectra of the collected electrolytes (DF: 100) were recorded to study the Pt concentration evolution with applied potential.

**Quantification of Pt-dissolution through UV-Vis spectroscopy:**

UV-Vis spectroscopy, a technique requiring low initial cost, may be used for quantification of the Pt-dissolution in chloride containing electrolytes from the catalyst samples. The electrochemical
dissolution of Pt in such baths takes place through formation of \([PtCl_4]^{2-}\) and/or \([PtCl_6]^{2-}\) complexes, which exhibit absorption peaks centered at ~265 nm and ~225 nm, respectively. The concentration of Pt in the electrolyte can be measured by measuring the corresponding absorbance, which is directly proportional to the concentration of absorbing species in solution (Beer Lambert law). Fig. 2 shows calibration curves made for both the complexes by using commercial Pt precursors: \(K_2PtCl_4\) solution in 0.1 M HCl for the \([PtCl_4]^{2-}\) complex and \(PtCl_4\) solution in 0.1 M HCl for the \([PtCl_6]^{2-}\) complex (\(PtCl_4 + 2HCl \rightarrow H_2PtCl_6\)).

![UV-Vis spectra](image)

Fig. 2: UV-Vis spectra of (a) the \(K_2PtCl_4\) solutions in 0.1 M HCl with varying Pt concentrations between 0.75 and 20 µg/mL and (b) the \(PtCl_4\) solutions in 0.1 M HCl with varying Pt concentrations between 0.155 and 2.5 µg/mL. The Pt concentrations have been shown in legends. Variations of the
area under the absorption peak with Pt concentrations corresponding to (c) the $[PtCl_4]^{2-}$ and (d) the $[PtCl_6]^{2-}$ complexes.

**Structural characterizations:** X-ray diffraction (XRD) patterns of the pristine and the electrochemically treated electrodes were collected using a Rigaku Miniflex 600 X-ray diffractometer (X-ray radiation: Cu Kα ($\lambda = 1.5418 \text{ Å}$)). The Pt (111) diffraction peak at diffraction angle ($2\theta$) = 39.8° (JCPDS# 65-2868) was used to determine the average crystallite size of Pt-nanoparticles ($L$) using Scherer’s formula ($L = 0.9\lambda/\beta Cos\theta$; where $\beta$ is the width ($2\Delta\theta$) of the diffraction peak, determined from the full width at half maximum (FWHM) of Lorentz distribution ($\beta = \text{FWHM} \times \pi/2$).

Finally, transmission electron microscope (TEM; JEOL 3000F) was used to observe the size distribution and the abundance of the Pt-nanoparticles in pristine and electrochemically treated electrodes. Samples for TEM imaging were prepared by dispersing the catalyst layer in ethanol ultrasonically and drop casting on a Cu grid.

3. Results and discussion

3.1 Reliability studies

The reliability of the dissolution protocol was carefully evaluated and is reported in Table 1. The procedure produces relative standard deviation of less than 4% and is implemented for rest of the studies on Pt electrochemical dissolution optimization and mechanism interpretation.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Dissolution percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>53.0</td>
</tr>
<tr>
<td>2</td>
<td>52.2</td>
</tr>
<tr>
<td>3</td>
<td>55.9</td>
</tr>
<tr>
<td>Avg.</td>
<td>53.7</td>
</tr>
<tr>
<td>Rstdev</td>
<td>3.6%</td>
</tr>
</tbody>
</table>
The dissolution protocol established in this work has several advantages. Firstly, in electrodes, catalyst powder is physically localized by polymeric binder, which provides an easy solution for solid–liquid separation; moreover, catalyst ink is supported by an electron conductive layer, which provides excellent current distribution to reach catalyst nanoparticles. These greatly simplify experimental procedures. Secondly, graphite is an easily accessible material with good conductivity and high corrosion resistance. In contrast, metallic connections may corrode and cannot guarantee stable contact, consequently degree of dissolution; moreover, metallic impurities may disturb the dissolution mechanism and lead to misinterpretation. Dissolution and residual element quantification are complementary to each other and provide cross-checking possibility; furthermore, the high sensitivity of the instruments allows for minimal material use for accurate quantitative analysis. XRF provides instant, high throughput and reliable metallic element detection. Table 2 shows a comparison of the measurements of the dissolved Pt amounts in 0.1 M HCl during different experiments through AAS and XRF techniques. The %deviation of the XRF measurements from the corresponding AAS measurements lies within ±5%. The good correlation between the values determined through XRF and AAS suggests high reliability of the XRF technique. Further, the correspondence also suggests no significant deposition of Pt on the counter and reference electrodes during the dissolution study.

**Table 2:** Comparison of the measurements of the dissolved Pt amounts in 0.1 M HCl during different experiments through AAS and XRF techniques

<table>
<thead>
<tr>
<th>Electrode</th>
<th>Total Pt loading (µg)</th>
<th>Vol. (mL)</th>
<th>AAS conc. (DF: 200)</th>
<th>%dissolution (XRF)</th>
<th>Dissolved Pt from XRF (µg)</th>
<th>Dissolved Pt (µg) from AAS</th>
<th>%deviation</th>
</tr>
</thead>
</table>
3.2 Electrochemical parameters (potential range)

A number of electrochemical parameters, such as the potential window, waveform of the potential cycle (triangular, square, sawtooth, etc.), scan rate, etc., may be varied during the electrochemical dissolution process. Among these, the potential window, which determines the overall dissolution mechanism, is of high significance. While significant redeposition of the dissolved Pt species may take place for low potentials, higher potentials may lead to electrolysis of water, chlorine evolution and carbon corrosion, and hence, both the upper and the lower potential limits determine the overall dissolution mechanism.

To observe the electrochemical reactions during potentiodynamic treatment, cyclic voltammograms recorded by subjecting a typical Pt/C WE used in the present study to a potential cycling in 0.1 M HCl at a scan rate of 10 mV/s between different potential ranges are shown in Fig. 3a. A scan rate slower than that used for the dissolution study is used to make the Faradaic peaks observable that are obscure otherwise due to large capacitive currents at a scan rate of 100 mV/s (Fig. S1; supporting information S1). For comparison, cyclic voltammogram in 1 M H₂SO₄ exhibiting the characteristic peaks corresponding to (i) H⁺ adsorption, (ii) hydrogen evolution, (iii) hydrogen oxidation reaction (HOR), (iv) H⁺ desorption, (v) Pt oxidation, (vi) oxygen evolution/carbon corrosion, (vii) Pt-Oxide
reduction, etc., \cite{45} is also provided in Fig. 3a. In 0.1 M HCl, the peaks corresponding to H\textsuperscript{+} adsorption, H\textsuperscript{+} desorption, hydrogen evolution and HOR become less clear probably due to Cl\textsuperscript{-} coverage of the Pt surface. Peak corresponding to Cl\textsuperscript{-} adsorption can be observed at \textasciitilde1.3 V during positive-going scan, while the onset of Cl\textsuperscript{-} desorption during the negative-going scan takes place at \textasciitilde1.0 V and proceeds till \textasciitilde0.5 V, along with the Pt-Oxide reduction. Further, large currents for potentials \textgreater1.4 V during positive-going scan correspond mainly to chlorine evolution along with the oxygen evolution/carbon corrosion., while the current peak at \textasciitilde1.3 V for negative-going potentials (\textgreater1.3 V) is attributed to the chlorine reduction (Cl\textsubscript{2} + 2e\textsuperscript{-} \rightarrow 2Cl\textsuperscript{-} (1.358 V)).\cite{46}

The effect of the lower potential limit on the dissolution rate (in terms of %dissolution from WE: 10 mm disc) of a triangular waveform was studied by keeping the upper potential limit fixed at 1.6 V and varying the lower potential limit between 1.0 and 0.0 V during potential cycling in 0.1 M HCl at a scan rate of 100 mV/s for 50 cycles. Similarly, the effect of the upper potential limit on the dissolution rate of a triangular waveform was studied by keeping the lower potential limit fixed at 0.4 V and varying the upper potential limit between 1.0 and 1.7 V during potential cycling in 0.1 M HCl at a scan rate of 100 mV/s for 50 cycles. As shown in Fig. S1 (supporting information), the shape of a voltammogram changes significantly with \(N\) due to factors such as (i) the effective Pt surface area, (ii) the concentration of dissolved Pt species, (iii) the Pt-nanoparticle/carbon support interaction, (iv) the wettability of the electrode due to possible carbon corrosion from the GDL and the catalyst support and change of ionomer interface. With varying upper/lower potentials the voltammogram shapes change due to occurrence of difference redox reactions including hydrogen evolution, chlorine evolution, water electrolysis, etc.

The effect of varying the upper and lower potential limits on the %dissolution has been shown in Fig. 3b. For the lower potential limit study (upper potential: 1.6 V), no significant Pt-dissolution takes place for the lower potential higher than 0.8 V, while the %dissolution in specified \(N\) value decreases
for the lower potential limits below 0.4 V. Hence, a lower potential limit of 0.4 V is found optimal for the Pt recovery for the studied electrodes through electrochemical dissolution. On the other hand, %dissolution of Pt increases with increasing the upper potential limit from 1.0 to 1.7 V while keeping the lower potential limit fixed at 0.4 V. The observed monotonous increase of the %dissolution with increasing upper potential is in contrast to that reported by Kanamura et al. (2016), where a decrease of dissolution has been reported for potentials above 1.4 V during dissolution of Pt in 1 M HCl through square-wave potential cycling between lower potential of 0.1 V and varying upper potential values. This may be due to comparatively less redeposition of the dissolved Pt species at 0.4 V in the present study as compared to that at 0.1 V for the earlier report.

Chlorine evolution at potentials >1.36 V may also affect the Pt dissolution differently for square and triangular waveforms as in the later case, Cl₂ concentrations decrease significantly through its reduction and diffusion before the reduction of Pt-Oxides. However, to avoid significant chlorine evolution/carbon corrosion and to obtain a high dissolution rate, the upper potential limit in the present study has been restricted to 1.6 V.

Apart from optimization of the working potential window, the upper/lower potential limit studies shed light on the overall dissolution mechanism. The dissolution takes place by chloride complexing through equations (2) and (3) for potentials >0.75 V, along with parallel formation of Pt-Oxide layer through equations (4), (5) and (6). During the cathodic sweep, Pt-dissolution takes place through dissolution/chloride complexing of the Pt-Oxide layer, leading to a fresh Pt surface for potentials ranging between 1.0 and 0.4 V. The potential value sufficient to dissolve the Pt-oxide layer completely may be varied by varying the reaction kinetics by changing the parameters such as scan rate, Pt-nanoparticle/support interface, etc. The local potential at the nanoparticle surface may differ significantly from the externally applied potential due to the insulating property of the proton conductive polymer and the ionomer (thin film) encapsulating catalyst interface structure,
there is an increased electron transport resistance at the catalyst/support/ionomer interface.\textsuperscript{[49, 51]}

Therefore, the values reported in the study is specific for the PEMFC electrodes; nevertheless, the relative value and changing trend is representative and can be used for general understanding. Further, as the cathodic peaks corresponding to Cl\textsuperscript{-} desorption/Pt-Oxide reduction during negative-going scan in 0.1 M HCl (Fig. 3a) shift towards negative potentials with increasing the upper potential limit, the potential range for dissolution may be optimized further through choosing a smaller potential step (e.g. 100 or 50 mV) for the lower/upper potential limit study, instead of a potential step of 200 mV used in the present study.

\begin{equation}
[\text{PtCl}_4]^{2-} + 2e^- \rightarrow Pt + 4Cl^- (E^o = 0.75 \text{ V}) \tag{2}
\end{equation}

\begin{equation}
[\text{PtCl}_6]^{2-} + 2e^- \rightarrow [\text{PtCl}_4]^{2-} + 2Cl^- (E^o = 0.68 \text{ V}) \tag{3}
\end{equation}

\begin{equation}
P tO + 2H^+ + 2e^- \rightarrow Pt + H_2O (E^o = 0.88 \text{ V}) \tag{4}
\end{equation}

\begin{equation}
P tO_2 + 2H^+ + 2e^- \rightarrow PtO + H_2O (E^o = 1.01 \text{ V}) \tag{5}
\end{equation}

\begin{equation}
P tO_2 + 4H^+ + 4e^- \rightarrow Pt + 2H_2O (E^o = 1.00 \text{ V}) \tag{6}
\end{equation}

Further, redeposition of the dissolved Pt on the source Pt-nanoparticles or on the WE surface may take place through equations (2) and (3) for potentials below the Nernst potential.

Nernst equations corresponding to eq. (2) and (3) can be written as equations (7) and (8), respectively,

\begin{equation}
E = E^o - \frac{RT}{nF} \ln \left( \frac{a_{Cl^-}}{a_{[PtCl_4]^{2-}}} \right) \tag{7}
\end{equation}

\begin{equation}
E = E^o - \frac{RT}{nF} \ln \left( \frac{a_{[PtCl_4]^{2-}} \times a_{Cl^-}^2}{a_{[PtCl_6]^{2-}}} \right) \tag{8}
\end{equation}

where $E$ and $E^o$ are the Nernst and the standard redox potentials, R is the gas constant, T is the Temperature (K), F is the Faraday constant and $n$ is the number of electrons in half-reaction and $a_x$ is the activity of the species $x$.

For estimation of the Nernst potentials corresponding to dissolution or redeposition of Pt, the equilibrium $[\text{PtCl}_4]^{2-}$ concentration is assumed to be 5% of the $[\text{PtCl}_6]^{2-}$ concentration,
considering the fact that no significant fraction of $[PtCl_4]^{2-}$ is observed in the electrolyte. For typical Pt concentrations of Table 2 ($\leq 0.5$ mM) and Cl$^-$ concentration of 0.1 M, in the present study, the Nernst potentials corresponding to eq. (1) and (2) are estimated to be 0.73 and 0.78 V, respectively. However, the effect of redeposition on the %dissolution is observable for the lower potential limits $<0.4$ V (Fig. 3b). This may be attributed to combined effects of the increasing %dissolution through complete dissolution of Pt-Oxide and the reaction overpotential for Pt redeposition shifting the redeposition potential towards lower values.
Fig. 3: Cyclic voltammograms in 0.1 M HCl or 1 M H₂SO₄ at a scan rate of 10 mV/s. Peaks corresponding to various Faradaic reaction are labeled as- (i) H⁺ adsorption, (ii) hydrogen evolution, (iii) HOR, (iv) H⁺ desorption, (v) Pt oxidation, (vi) oxygen evolution/carbon corrosion, (vii) Pt-Oxide reduction, (viii) Cl⁻ adsorption, (ix) chlorine evolution, (x) chlorine reduction and (ix)
Cl⁻ desorption. (b) Variations of %dissolution of Pt from WE with the lower potential limit and the higher potential limits during potential cycling.

3.3 Effects of H⁺ and Cl⁻ concentrations

Electrolytes without Cl⁻ as complexing agents: In absence of Cl⁻ as complexing agent, no significant dissolution of Pt takes place. As an example, potential cycling of the PEMFC electrodes (10 mm diameter) between 0.4 and 1.6 V in 0.1 M, 1 M and 4 M nitric acid for 100 cycles at a scan rate of 100 mV/s exhibits %dissolutions of ~0.3%, 1.8% and 3.6%, respectively. Moreover, increasing the \( N \) does not seem to increase the %dissolution proportionally. For example, 2500 potential cycles (0.4 and 1.6 V; scan rate: 100 mV/s) in 0.1 M HNO₃ exhibit a %dissolution of 1.2%. Further, increasing the NO₃⁻ concentration by adding NH₄NO₃ does not improve the dissolution significantly (supporting information: S2). Similar observations of low %dissolution of Pt in other Cl⁻ free electrolytes such as 1 M H₂SO₄ and 0.1 M HClO₄ have also been made in our previous studies.[31, 52-53] Hence, in absence of complexing agents, very little amount of Pt was removed from the WE due to the unstable nature of the Pt-Oxides formed at potentials higher than ~1 V at the low pH values of the studied electrolytes[54] and high tendency for redeposition. However, the average Pt-nanoparticle size increases during the electrochemical treatment due to Ostwald ripening.[31, 53] For example, the crystallite size of the electrode subjected to potential cycling (0.4-1.6 V; 100 mV/s) for 200 cycles in 1 M HNO₃ is observed to be 3.1 nm as compared to 2.0 nm of the pristine electrode (supporting information: S3).

Electrolytes with Cl⁻ as complexing agents: Dissolution of Pt in presence of Cl⁻ may take place at comparatively lower potentials through equations (2) and (3). The potential vs. pH diagram in presence of Cl⁻ suggests the dissolution potential to be independent for a broad range of pH between 0 and ~6.[41] Fig. 4a shows a comparison of the %dissolution values in 0.1 M and 1 M HCl electrolytes,
exhibiting a clear enhancement of dissolution rate with the HCl concentration. The most efficient Pt-dissolution in 1 M HCl, reaching to ~95% dissolution in 50 cycles may be compared with that in 0.1 M HCl, where 250 potential cycles produced ~90% dissolution. However, the first derivative \( (d(%\text{dissolution})/dN; \%\text{dissolution/cycle}) \) decreases sharply with increasing \( N \) due to the decreasing fraction of undissolved small Pt nanoparticles, higher Pt concentration in the electrolyte and therefore higher probability of redeposition. The estimated average rate of Pt-dissolution/cycle for the first two cycles in 1 M HCl is ~30 µg/cycle.

Fig. 4: Variations of %dissolution and %dissolution/cycle vs. number of cycles for electrochemical dissolution in (a) 0.1M, 1 M HCl and (b) 0.5 M NH₄Cl electrolytes. Variation of %dissolution during electrochemical dissolution in different electrolytes through potential cycling (potential
range: 0.4-1.6V; scan rate: 100 mV/s) for 50 cycles with (c) varying the NH4\(^+\) concentration (Cl\(^-\) concentration: 0.5 M) and (d) varying both the NH4\(^+\) and the Cl\(^-\) concentrations.

**Pt-dissolution in acid-free baths:** Apart from dilute acidic baths i.e. 0.1 M HCl, Pt dissolution in acid-free baths such as aqueous solutions of NH4Cl or NaCl has been demonstrated. Electrochemical dissolution studies in the NH4Cl or NaCl baths have been performed through potential cycling between 0.4 and 1.6 V at a scan rate of 100 mV/s for a specified value of \(N\). Fig.4b shows the variations of %dissolution and d(%dissolution)/d\(N\) with \(N\) in 0.5 M NH4Cl. With ~70% dissolution in the first 200 cycles, the acid-free electrolyte exhibits promising performance. Furthermore, Fig. 4c exhibits the impact of the NH4\(^+\) concentration (Cl\(^-\) concentration: 0.5 M; adjusted by NaCl) on the %dissolution of Pt during potential cycling for 50 cycles between 0.4 and 1.6 V at a scan rate of 100 mV/s. As shown in Fig. 4d, potential cycling (0.4-1.6 V; 100 mV/s; 50 cycles) in a 0.5 M NH4Cl bath exhibits 5 times higher %dissolution of Pt as compared to that in a 0.5 M NaCl bath. This is due to the fact that the dissolution potentials remain constant for pH below ~6 in Cl\(^-\) containing electrolytes, and the dissolution takes place through equations (2) and (3). However, for pH >6, the dissolution takes place through formation of Pt(OH)\(_2\) rather than through the [PtCl\(_{x}\)]\(^{2-}\) complexes.\[^{41}\]

UV-vis spectrum of the dissolved Pt species in 0.5 M NaCl (pH 7) through 50 cycles between 0.4 and 1.6 V at a scan rate of 100 mV/s is shown in Fig. S3 (supporting information S4). The spectrum does not show peaks corresponding to either of the [PtCl\(_{4}\)]\(^{2-}\) or [PtCl\(_{6}\)]\(^{2-}\) complexes and therefore the %dissolution remains comparable to that observed for the Cl\(^-\) free electrolytes.

Further, for the Cl\(^-\) concentrations of 1 M and in presence of NH4Cl higher NH4\(^+\) concentration leads to higher %dissolution. This may be attributed to (i) the relatively higher acidic nature of the 0.5 M NH4Cl electrolyte (pH: ~4.5) as compared to the neutral NaCl electrolyte leading to dissolution
through formation of the $[\text{PtCl}_x]^{2-}$ complexes and (ii) the enhanced reaction kinetics due to lower solubility of the dissolution product $(\text{NH}_4)_2\text{PtCl}_6$.

**Effect of electrolyte convection flow:** The potential limit study suggests redeposition of the dissolved Pt species, especially for potentials below 0.5 V during the electrochemical potential cycling. Using identical location scanning electron microscope (IL-SEM) imaging of the electrochemically degraded electrodes, Hodnik et al. (2013) have observed a three-dimensional Ostwald ripening by the Pt dissolution from inner layers and redeposition of the dissolved Pt species on the top layers of a thin-film catalyst. Hence, the rate of redeposition of the dissolved Pt species is expected to increase with increasing their local concentration in proximity of the electrode.

Comparison of %dissolution values for Pt dissolution through potential cycling between 0.4 and 1.6 V at a scan rate of 10 mV/s in 0.1 M HCl for 25 cycles with or without stirring the electrolyte suggests large enhancement of the dissolution rate. Under identical conditions, the %dissolution increases form ~16% for the stagnant electrolyte to ~80% for the electrolyte stirred at 750 rpm. This large enhancement of %dissolution may be attributed to the fact that in absence of convection flow, concentrations of the dissolved Pt species in proximity of the Pt nanoparticles remain high in the stagnant electrolyte as compared to those in the stirred electrolyte, leading to high rate of redeposition of the dissolved Pt species and hence lower % dissolution.

**3.6 Electrode interface influence on Pt catalyst dissolution**

Most PEMFC electrodes consist of a mixture of the carbon supported Pt-nanoparticles and the proton conducting ionomer (e.g. Nafion®) coated on the GDL. The catalyst/ionomer interface has been reported to affect the electrochemical catalytic performance of the electrode significantly. The catalyst/ionomer interface may similarly affect the Pt/electrolyte interaction, and hence the Pt-dissolution during the electrochemical treatment. To study the impact of catalyst/ionomer interface on the Pt-dissolution, PEMFC electrodes having different catalyst ink compositions with Nafion®
contents of 0, 15, 30 and 45 wt.% were studied. Potentiodynamic dissolution through potential cycling (potential range: 0.4-1.6V; scan rate: 100 mV/s) for 25 cycles in 0.1M HCl shows different %dissolution values with different ionomer contents (Fig. 5a). This series of electrodes are more prone to platinum dissolution than the electrodes used for the other experiments of this study, which might be due to their more open electrode structure created by the spray coating method rather than the tape casting used for the later ones. High %dissolution for the electrode with 0% Nafion® content as compared to those with higher Nafion® contents suggests the significant hindrance of the Nafion® on the dissolution process. This may be attributed to the low diffusion coefficients or repulsion from negative charges of the \([PtCl_4]^{2−}\) and \([PtCl_6]^{2−}\) complexes in the cation exchange resin Nafion® as compared to those in the aqueous electrolyte. This may increase local concentrations of the dissolved Pt species and hence lead to higher redeposition. Additionally, formation of Pt nanoparticles inside the ionomer through reduction of the Pt species migrated inside the ionomer membrane, a mechanism similar to that observed during electrode degradation in PEMFCs by reduction of the Pt species through the crossed-over hydrogen,[60-61] might play a role. At the electronically connected carbon/ionomer interface, reduction of the entrapped Pt species during negative-going potential scan may take place.

Further, the slightly increasing Pt dissolution trend with increasing ionomer content (15, 30 and 45 wt.%) might be attributed to the more accessible Pt surface due to ionomer agglomeration at high concentration.[56] XRD patterns of the pristine electrodes (Fig. 5b) with different Nafion® content show similar peak widths for the Pt (111) peak, eliminating the possibility of initially different particle sizes. The XRD patterns of the post-dissolution electrodes show a marginal increase of the particle sizes (−5%).
Fig. 5: (a) Variation of %dissolution for electrochemical dissolution in 0.1M HCl through potential cycling (potential range: 0.4-1.6V; scan rate: 100 mV/s) for 25 cycles. (b) XRD patterns of pristine and post-dissolution electrodes with different Nafion® contents, showing the Pt (111) and Pt (200) peaks at 2θ values ~39.9° and 46.1°, respectively. The inset of (b) shows the XRD patterns for a broader scan range.

3.4 Dissolution product and electrode structure evolution during dissolution

Recovery of spent catalysts through electrochemical dissolution of Pt may be achieved by using the electrolytes containing the species that form stable Pt complexes. Electrochemical dissolution of Pt in presence of Cl- leads to formation of Pt complexes. As can be seen from the UV-Vis spectra of the post-dissolution electrolytes. Fig. 6a shows the UV-vis spectra of the post-dissolution electrolytes for electrochemical Pt-dissolution studies by potential cycling in (i) 0.1 M HCl (25 cycles) and (ii) 0.5 M NH₄Cl (50 cycles) between 0.4 and 1.6 V at a scan rate of 100 mV/s. Here, the samples were diluted appropriately using 0.1 M HCl or 0.1 M NH₄Cl before the UV-Vis measurements to obtain comparable absorbance values below 0.5. For comparison, UV-Vis spectra of commercial Pt precursors, namely H₂PtCl₆ and K₂PtCl₄ solutions in 0.1 M HCl exhibiting absorption peaks corresponding respectively to the [PtCl₆]²⁻ (265 nm) and the [PtCl₄]²⁻ (225 nm) complexes are
also included. The UV-Vis spectra of the post-dissolution electrolytes show characteristics similar to that of the H2PtCl6. This confirms the chemical state of the dissolved Pt to be $[PtCl_6]^{2-}$ complex for both of the electrolytes. To illustrate the reaction process, fig. 6b shows the UV-vis spectra of the electrolyte 0.1 M HCl at different potentials during the first potential cycle with start, maximum, minimum and end potential values of 1.2 V, 1.6 V, 0.4 V and 1.6 V, respectively. At the beginning of potential cycling from 1.2 V to positive-going segment (1.2–1.6 V), Pt dissolution takes place through formation of $[PtCl_4]^{2-}$ species. During the negative-going scan from 1.6 to 0.8 V, concentration of the $[PtCl_4]^{2-}$ species increases, while it decreases for the scan from 0.8 to 0.4 V. This decrease in the $[PtCl_4]^{2-}$ species concentration may be attributed to their redeposition as Pt0.[24, 31] Further, the concentration of $[PtCl_6]^{2-}$ species appears for the first time during the negative-going scan from 0.8 to 0.6 V. The appearance of the absorption peak corresponding to $[PtCl_6]^{2-}$ complex may be attributed to the Pt-dissolution through the Pt-Oxide removal below 1.0 V during negative-going scan, as has been observed in the corresponding cyclic voltammogram of Fig. 3a. Here, the fact that the sample collected at a certain potential for the UV-vis measurement may not represent the instantaneous concentration of the Pt species due to a time lag between their production and uniform distribution in the electrolyte. This may explain the first appearance of $[PtCl_6]^{2-}$ species during the 0.8–0.6 segment, despite their formation during the 1.0–0.8 segment of the negative-going scan. Further, the nearly constant concentration of $[PtCl_6]^{2-}$ species during the negative-going (0.6–0.4 V) and the positive-going (0.4-0.8 V) scans suggests no significant dissolution or redeposition of Pt taking place in this potential widow. The $[PtCl_6]^{2-}$ concentration increases for the positive –going segment between 0.8 and 1.6 V due to anodic dissolution of Pt through formation of Pt-oxides.[20] Here, the significant variation of the background below 220 nm, possibly due to change in the Cl- ion concentration, however makes it difficult to comment on the $[PtCl_6]^{2-}$ concentration. Nevertheless, unlike the first positive going segment between 1.2 and 1.6 V, increase of the $[PtCl_6]^{2-}$ concentration
for the corresponding second positive going segment suggests different Pt dissolution mechanisms for the two cases, possibly due to the presence of intrinsic Pt-Oxide in the former case. Further investigations may be performed to understand the mechanisms. Also, real-time monitoring of the UV-vis spectra through flow-cell may be employed to observe the evolution of $\text{[PtCl}_4^2-]/\text{[PtCl}_6^2-$ ratio during the potentiodynamic dissolution in presence of Cl$^-$. 

Fig. 6: (a) UV-Vis spectra of different post-dissolution electrolytes (potentiodynamic dissolution of Pt from the PEMFC electrode by potential cycling in 0.1 M HCl (25 cycles) and 0.5 M NH$_4$Cl (50 cycles) between 0.4 and 1.6 V at a scan rate of 100 mV/s. For comparison, UV-Vis spectra of H$_2$PtCl$_6$ and K$_2$PtCl$_4$ solutions in 0.1 M HCl are also included. (b) UV-vis spectra of the 0.1 M HCl electrolyte samples collected at different potentials during the first potential cycle. Legends show the additional potential sweep segment completed between the consecutive electrolyte sample collections.

Further, XRD patterns of the electrochemically treated post-dissolution electrodes treated by potential cycling between 0.4 and 1.6 V at a scan rate of 100 mV/s in 0.1 M HCl (25 cycles) and 1 in 1 M NH$_4$Cl (200 cycles) electrodes are shown in Fig. 7a and 6b, respectively. For comparison, XRD pattern of a pristine electrode is also included. The XRD pattern of pristine and HCl treated electrodes
show diffraction peaks corresponding to both the graphitic carbon from GDL and the Pt-nanoparticles.[31, 52] As shown in the inset of Fig. 7a, the decreased peak width of the diffraction for the post-dissolution sample suggests an increased average size of crystallites from 2.0 nm for pristine electrode to 2.4 nm for the post-dissolution electrode (~25% dissolution). The crystallite growth may be attributed to the faster dissolution of the smaller particles and/or the growth of larger particles (Ostwald ripening). Further, the XRD patterns of the pristine and the post-dissolution electrode treated in 1 M NH₄Cl are shown in Fig. 7b. The ‘extra’ diffraction peaks present in the pattern corresponding to the post-dissolution electrode suggest the presence of (NH₄)₂PtCl₆ deposits on the electrode during dissolution process, owing to its low solubility in the 1 M NH₄Cl electrolyte. Hence, for the NH₄Cl electrolyte, the dissolution product appears both in form of the (NH₄)₂PtCl₆ deposits[62] on electrode and the [PtCl₆]²⁻ complex in the solution. On the other hand, the post-dissolution electrode treated in 0.1 HCl shows no ‘extra’ peaks compared to the pristine electrode, suggesting the undissolved Pt to be in form of Pt-nanoparticles and hence the dissolution product to be in form of [PtCl₆]²⁻ complex in the solution.

Fig. 7: XRD patterns of the electrochemically treated post-dissolution electrodes treated by potential cycling between 0.4 and 1.6 V at a scan rate of 100 mV/s in 0.1 M HCl (25 cycles) and
in 1 M NH₄Cl (200 cycles) electrodes. For comparison, XRD pattern of pristine electrode is also included. Diffraction peaks corresponding to (NH₄)₂PtCl₆ have been marked with (*).

Comparison between the TEM images of the pristine (fig. 8a) and the post-dissolution study electrodes (fig. 8b-8d) reveals significant reduction of Pt particle density during dissolution. However, different TEM images of the later reveal inhomogeneous distribution of the Pt-nanoparticles in different areas of the electrode. The inhomogeneous dissolution could be due to the finite thickness of the catalyst layer where the top layer is exposed to the electrolyte most effectively while the inner layer of the catalyst layer is partially obscured. Significant growth of the particles, can be seen in Fig. 8b-8d, with larger particle size for smaller number density. As the XRD patterns suggest growth of crystallites with dissolution treatment, the growth of particle size observed from the TEM images may be attributed mainly to the Ostwald ripening (redeposition of the dissolved Pt species on the source Pt nanoparticles) along with other mechanisms such as particle agglomeration. Here, the redeposition probably takes place on the source Pt nanoparticles rather than formation of new particles on the carbon support. However, cross-sectional imaging of the post-dissolution electrodes may be performed to understand the redeposition process during the electrochemical dissolution.[55] The dissolution efficiency may be enhanced significantly by using strategies to reduce the redeposition of dissolved Pt species.
Fig. 8: TEM images of (a) the pristine and (b-d) the post-dissolution study (60% dissolution) electrodes.

**Upscaling for end-of-life MEA**

The presented electrochemical dissolution route is being explored towards its feasibility for the recovery of Pt from end-of-life MEAs. The initial challenge towards recycling Pt from the end-of-life MEAs is the delamination of the MEAs, necessary to make the catalyst layer accessible to the
electrolyte. During delamination, the catalyst layers attach partially to the PEM, making it difficult to provide electronic connectivity for the potentiodynamic dissolution. Strategies addressing this issue may include the delamination with preferred attachment of the catalyst layer to the GDL, the separation of the catalyst from the PEM followed by the dissolution of Pt, etc. Moreover, % Pt dissolution from different parts of a large electrode (e.g. 10 × 10 cm²) varies significantly depending on the distance from the connection to the external circuit (low dissolution for areas far from the connection point). However, the issue can be solved by providing connections at multiple points. To summarize, the presented process to recover Pt electrocatalysts through potentiodynamic dissolution may be scaled up after solving such practical issues.

4. Conclusions
In conclusion, efficient recycling of precious metal catalysts supported on conductive substrates through potentiodynamic treatment under mild conditions has been studied. High activity of the nanoparticulate catalysts helps their faster dissolution in dilute acidic/acid-free chloride baths. The dissolution rate is influenced significantly by parameters such as the potential window, electrolyte composition, electrode interface, etc. A potential range between 0.4 and 1.6 V was found optimal for the typical PEMFC electrodes under study. Poor acidic and acid free Cl⁻ containing electrolytes such as 0.1-1.0 M HCl and 0.5-1.0 M NH₄Cl show efficient Pt recovery. For an initial Pt loading of ~470 µg, an average rate of Pt-dissolution/cycle as high as ~30 µg/cycle is achieved during potential cycling between 0.4 and 1.6 V in (scan rate: 100 mV/s) in 1.0 M HCl. In summary, the electrochemical dissolution route to recycle platinum catalyst from energy conversation systems presents an important recycling process and significant advance to sustain the green technologies.

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References


Supporting information on **Sustainable platinum recycling through electrochemical dissolution of platinum nanoparticles from fuel cell electrodes**

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**S1: Cyclic voltammograms of dissolution study (potential limit)**

![Cyclic voltammograms for (a) lower and (b) upper potential limit studies in 0.1 M HCl for 50 cycles at a scan rate of 100 mV/s](image)

Fig. S1: Cyclic voltammograms for (a) lower and (b) upper potential limit studies in 0.1 M HCl for 50 cycles at a scan rate of 100 mV/s

**S2: Dissolution of Pt in nitric acid based electrolytes**

Table S1 shows the values of %dissolution of Pt in various nitric acid/based electrolytes during potential cycling between 0.4 and 1.6 V at a scan rate of 100 mV/s for varying number of potential cycles.
Table S1: %dissolution of Pt during electrochemical treatment in nitric acid

<table>
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<tr>
<th>Electrolyte</th>
<th>Number of cycles</th>
<th>%Dissolution</th>
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</thead>
<tbody>
<tr>
<td>0.1 M HNO₃</td>
<td>100</td>
<td>0.3</td>
</tr>
<tr>
<td>0.1 M HNO₃</td>
<td>200</td>
<td>0.6</td>
</tr>
<tr>
<td>0.1 M HNO₃</td>
<td>2500</td>
<td>1.2</td>
</tr>
<tr>
<td>0.1 M HNO₃ + 0.1 M NH₄NO₃</td>
<td>100</td>
<td>1.6</td>
</tr>
<tr>
<td>0.1 M HNO₃ + 0.1 M NH₄NO₃</td>
<td>200</td>
<td>3.6</td>
</tr>
<tr>
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<td>6.8</td>
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</tr>
<tr>
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</tr>
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S3: Particle growth in HNO₃ electrolyte

XRD patterns of the post-dissolution electrode treated by potentiala cycling between 0.4 and 1.6 V at a scan rate of 100 mV/s in 1 M 1 M HNO₃ (200 cycles) and the pristine electrode are shown in Fig. S2. Despite a %dissolution of 3.3%, the post dissolution electrode shows significant growth of crystallite size, as revealed by the decreased peak width of the Pt(111) peak at 20 value of 39.9°.
Fig. S2: XRD patterns of the post-dissolution electrode treated by potential cycling between 0.4 and 1.6 V at a scan rate of 100 mV/s in 1 M HNO₃ (200 cycles) and the pristine electrode

S4: UV-vis spectrum of Pt species dissolved in 0.5 M NaCl

Fig. S3: UV-Vis spectrum of Pt species dissolved in 0.5 M NaCl through potential cycling between 0.4 and 1.6 V (50 cycles) at a scan rate of 100 mV/s. For reference, background spectrum of 0.5 M NaCl is also shown. Spectra recorded w.r.t. 0.5 M NaCl reference.